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Molecular Dynamics Simulation of Nucleation Process of SWNT from a Metal Particle on a Substrate YASUSHI SHIBUTA, SHIGEO MARUYAMA, Dept. of Mech. Eng., The University of Tokyo — Nucleation process of single-walled carbon nanotubes (SWNTs) from a transition metal cluster on a substrate is studied using classical molecular dynamics (MD) simulations. For describing the effect of the substrate, averaged one-dimensional Lennard-Jones potential is employed between the metal cluster and the bottom boundary of the simulation cell. As the initial condition, a Ni500 cluster is placed on the bottom boundary of the cubic cell of 20 nm. The number of carbon atoms is adjusted to achieve the constant density by adding a new carbon atom to the cell when the metal cluster dissolves a carbon atom. As the metal cluster dissolves carbon atoms, the cluster becomes more wetting to the substrate. This may be due to the different wettability between pure metal and metal-carbide. Graphite structure gradually precipitates from the edge of the cluster. Nucleation mechanism of SWNTs will be discussed by comparing with the simulation using the floated catalyst.

Prefer Oral Session
 Prefer Poster Session

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